Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

Claim 1 (currently amended): A prepackaged aqueous pharmaceutical composition <u>in</u> <u>liquid form</u> for the treatment of <u>treating a patient having at least one</u> cardiac condition[s], <u>the pharmaceutical composition</u> comprising:

(a) at least two different pharmacologically active agents a first pharmacologically active agent in a first dosage amount and at least a second pharmacologically agent in a second dosage amount for the treatment of [a] the at least one cardiac condition[;],

wherein the choice of the at least first and at least second pharmacologically active agents and the first and second dosage amounts is dependent upon the patient's characteristics;

- (b) a buffering agent to buffer said composition; and
- (c) an osmotic-adjusting agent[.],

the pharmaceutical composition being packaged in a plurality of separate dispensers, each of the dispensers comprising a single day's dose of the at least first and at least second pharmacologically active agents.

Claim 2 (currently amended): A method of forming a prepackaged aqueous pharmaceutical composition in liquid form for the treatment of at least one cardiac condition[s], said method comprising the steps of:

mixing at least <u>a first</u> two different pharmacologically active agent[s] <u>and at least a second pharmacologically active agent;</u>

adding a buffering agent to said mixed at least first two different pharmacologically active agent[s] and at least second pharmacologically active agent; and

adding an osmotic-adjusting agent to said mixed mixture of the buffering agent, the at least first pharmacologically active agent and the at least second two-different pharmacologically active agents[s.]; and

packaging the composition in a plurality of separate dispensers, each of the dispensers comprising a single day's dose of the at least first and at least second pharmacologically active agents.

Claim 3 (currently amended): A process for the administration of a prepackaged aqueous solution or dispersion of at least a first pharmacologically active agent and at least a second two different pharmacologically active agent[s] for treating a patient having at least one the treatment of a cardiac condition, comprising:

selecting a predetermined first dosage amount for the at least first pharmacologically active agent each of said and a second dosage amount for the at least two different second pharmacologically active agent[s], said first and second dosage amounts selected based upon the patient's patient characteristics;

mixing said at least first pharmacologically active agent at a first dosage amount and the at least second pharmacologically active agent at a second dosage amount[s] with a buffering agent and an osmotic-adjusting agent, said mixture of the buffering agent, the osmotic-adjusting agent, the at least first pharmacologically active agent[s] and the at least second

<u>pharmacologically active agent</u> having stability with one another <u>being stable</u> in aqueous solution; and

packaging the mixture in a plurality of separate dispensers, each of the dispensers

comprising a single day's dose of the at least first and at least second pharmacologically active

agents; and

providing said pharmacologically active agents having stability with one another in aqueous solution the packaged mixture to a suitable patient for oral administration.

Claim 4 (currently amended): The composition according to claim 1 wherein said at least two different first pharmacologically active agent and at least second pharmacologically active agents are selected from the group consisting of diuretics, cardiac glycocides glycosides, beta blockers, nitrates, antiplatelets, vitamins, nitroceuticals nutraceuticals, angiotensin converting enzyme inhibitors, and calcium channel elockers blockers.

Claim 5 (original): The composition according to claim 1 wherein said buffering agent comprises at least one of acetate, glutamate, citrate, tartrate, benzoate, lactate, gluconate, phosphate and glycine.

Claim 6 (original): The composition according to claim 1 wherein said osmotic-adjusting agent comprises at least one of sodium chloride, dextrose, sodium bicarbonate, calcium chloride, potassium chloride, sodium lactate, Ringer's solution and lactated Ringer's solution.

Claim 7 (currently amended): The <u>method of composition according to claim 2</u> wherein said at least <u>first pharmacologically active agent and at least second two different</u> pharmacologically active agents are selected from the group consisting of diuretics, cardiac <u>glycocides glycosides</u>, beta blockers, nitrates, antiplatelets, vitamins, <u>nitroceuticals</u> <u>nutraceuticals</u>, angiotensin converting enzyme inhibitors, and calcium channel blockers.

Claim 8 (currently amended): The <u>method of emposition according to claim 2</u> wherein said buffering agent comprises at least one of acetate, glutamate, citrate, tartrate, benzoate, lactate, gluconate, phosphate and glycine.

Claim 9 (currently amended): The <u>method of eomposition according to</u> claim 2 wherein said osmotic-adjusting agent comprises at least one of sodium chloride, dextrose, sodium bicarbonate, calcium chloride, potassium chloride, sodium lactate, Ringer's solution and lactated Ringer's solution.

Claim 10 (currently amended): The eomposition according to process of claim 3 wherein said at least two different first pharmacologically active agent and at least second pharmacologically active agents are selected from the group consisting of diuretics, cardiac glycosides, beta blockers, nitrates, antiplatelets, vitamins, nitroceuticals nutraceuticals, angiotensin converting enzyme inhibitors, and calcium channel elockers blockers.

Claim 11 (currently amended): The eomposition according to process of claim 3 wherein said buffering agent comprises at least one of acetate, glutamate, citrate, tartrate, benzoate, lactate, gluconate, phosphate and glycine.

Claim 12 (currently amended): The eomposition according to process of claim 3 wherein said osmotic-adjusting agent comprises at least one of sodium chloride, dextrose, sodium bicarbonate, calcium chloride, potassium chloride, sodium lactate, Ringer's solution and lactated Ringer's solution.

Claim 13 (new): The prepackaged pharmaceutical composition of claim 1, wherein each dispenser of the plurality of dispensers comprises a twist-off cap.